Serum IgE antibodies to *Trichophyton* in patients with urticaria, angioedema, asthma, and rhinitis: Development of a radioallergosorbent test


A series of patients was identified who demonstrated immediate positive skin tests to intradermal *Trichophyton* extract. These skin responses did not correlate with other fungal skin tests and were present both in atopic and nonatopic patients. The individuals demonstrating positive immediate skin tests included patients with urticaria, angioedema, asthma, and/or rhinitis, as well as five of 34 normal control subjects. Most skin test positive individuals had a history of local fungal infection and clinical signs suggestive of fungal infection. By use of *Trichophyton* tonsurans extract linked to Sepharose as the immunosorbent, it was possible to measure IgE antibodies in 26/30 sera from skin test positive individuals. With strongly positive sera, RAST bound up to 30% of the radiolabeled anti-IgE added. The results confirm that most skin test positive individuals have IgE-mediated hypersensitivity to *Trichophyton*. These observations support the older view that absorption of dermatophyte allergen through the skin should be considered as a possible cause of allergic disease. (J ALLERGY CLIN IMMUNOL 1987;79:40-5.)

The possible role of immediate hypersensitivity to *Trichophyton* allergens in diseases such as urticaria, rhinitis, and asthma was widely discussed during the 1930s. Two articles in the last two decades have suggested a role for *Trichophyton* antigens in chronic urticaria. It has also been reported that podiatrists become immune to *Trichophyton* and develop symptoms on exposure to the dust from infected toenails.

There have been several in vitro studies on the immune response to *Trichophyton* antigens. These studies have suggested that the nature of the immune response and the atopic state of the individual can alter the clinical course of the fungal infection but have not related these infections to any other disease. Surprisingly, there have been very few articles on the in vitro measurement of IgE antibody to *Trichophyton*. However, it has been reported that IgE antibody to fungal antigens can be measured in the serum of patients from New Guinea who have severe tinea imbracata. Those results were obtained by use of an ELISA assay on microtiter plates. There has also been a study from Denmark demonstrating that IgE antibody to *T. rubrum* can be detected by use of CRIE. However, a RAST is not currently available, and there are no studies of measurements of IgE antibody to *Trichophyton* in sera from patients with allergic diseases and fungal infection.

We became interested in immediate hypersensitivity to *Trichophyton* because we found several patients who had urticaria or recurrent angioedema associated with positive immediate skin tests to intradermal *Trichophyton* and who also had signs of fungal infection of their nails, feet, and/or groins. Further skin testing revealed positive skin tests in patients with asthma, some of whom had no other positive skin tests and also had signs of fungal infection. We report here the development of a RAST with *T. tonsurans* extract.

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**Abbreviations used**

- BBS: Borate-buffered saline, pH 8.0
- CRIE: Crossed radioimmunoelectrophoresis
- SAS: Saturated ammonium sulfate
- OD: Optical density
linked to Sepharose as the immunosorbent. The results demonstrate that IgE antibody to *Trichophyton* can be detected in serum from most individuals with strong positive immediate skin tests. These results are not restricted to patients with urticaria or asthma, since we have also found positive results among normal control subjects. Nonetheless, we believe our results suggest that the role of *Trichophyton* as an "intrinsic" allergen should be reconsidered.

**PATIENTS AND METHODS**

Sera were obtained from patients evaluated in the University of Virginia Allergy Clinic, including 27 patients with urticaria or recurrent angioedema, 39 patients with asthma, and seven patients with rhinitis. The patients with asthma included 15 patients who had demonstrated strongly positive skin tests to *Trichophyton* and 21 who had negative or weakly positive skin tests to *Trichophyton*. The patients with urticaria or recurrent angioedema included eight with strongly positive skin tests and 19 patients with negative or weakly positive skin tests to *Trichophyton*. Two of the seven patients with rhinitis had positive skin tests to *Trichophyton*. The patients with negative skin tests were selected to include patients with strong positive skin tests to other allergens. Thirty-four volunteers who were employed in the hospital were skin tested with both mite and *Trichophyton* extracts, and evaluated in the same way.

**Extracts and skin testing**

Skin testing was carried out with a mixture of *T. tonsurans*, *T. rubrum*, and *T. mentagrophytes* 1:10 w/v (Hollister-Stier, Spokane, Wash.). This extract was used undiluted for prick tests and 0.02 ml of 1:20 dilution (i.e., 1:200 w/v) was used for intradermal skin testing. Skin tests were graded 2 + = >5 by 5 mm wheal on prick test or >8 by 8 mm on intradermal tests and 1 + = >5 by 5 mm wheal on intradermal skin test. Patients were also skin tested with extracts of *Alternaria*, *Cladosporium*, and *Aspergillus*, as well as a range of other inhalant allergens as part of their routine evaluation. Skin tests to these three fungi were carried out by use of prick test and intradermal technique, as with *Trichophyton* and evaluated in the same way.

Cultures of *T. tonsurans* and *T. rubrum* that had been filtered from the fluid medium, rinsed, and freeze-dried were kindly provided by Hollister-Stier. Cultures were extracted with BBS, pH 8.0, at 1:7 w/v overnight in the cold, centrifuged to remove debris, and dialyzed. The mixed *Trichophyton* extract was also dialyzed against BBS. Extracts were compared according to their OD at 280 nm; 1 ml of extract with an OD of 1.0 equaled 1 OD unit. In some experiments extracts were fractionated with SAS by precipitation with 60% SAS followed by precipitation at 90%. SAS precipitate were dialyzed against three changes of BBS. Sepharose was then washed three times with bicarbonate buffer, incubated with 1 mol/L of ethanolamine for 2 hours, and washed with acetate buffer, pH 4.0, three times and with BBS three times. Sepharose was suspended to a final volume of 9.0 ml/gr of dry gel.

**TABLE I. Effects of species of *Trichophyton*, partial fractionation of extracts, and immunosorbent on RAST for *Trichophyton*-specific IgE antibodies**

<table>
<thead>
<tr>
<th>Serum</th>
<th><strong>Cellulose discs</strong></th>
<th><strong>Sepharose</strong></th>
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</thead>
<tbody>
<tr>
<td></td>
<td><em>T. tonsurans</em></td>
<td><em>T. rubrum</em></td>
</tr>
<tr>
<td></td>
<td>Whole extract</td>
<td>60% to 90% SAS</td>
</tr>
<tr>
<td>HD</td>
<td>9.1</td>
<td>11.8</td>
</tr>
<tr>
<td>RJ</td>
<td>1.8</td>
<td>2.6</td>
</tr>
<tr>
<td>SS</td>
<td>1.5</td>
<td>1.2</td>
</tr>
<tr>
<td>Controls</td>
<td>0.63</td>
<td>0.66</td>
</tr>
</tbody>
</table>

*Values are the mean of duplicate values for percent radioactive counts bound. All sera were assayed undiluted with 0.1 ml. For all other reagents in the assay, 50% horse serum was used as diluent. In separate experiments with 1% bovine serum albumin as a diluent, results for the same four skin test negative control subjects varied from 1.3% to 2.6% of counts bound. Extracts and 60% to 90% SAS fractions were dialyzed for 2 days against four changes of BBS.

†Mixed extract containing *T. tonsurans*, *T. rubrum*, and *T. mentagrophytes*.

*In a separate experiment, values obtained with 60% to 90% SAS fraction on Sepharose were very similar.*
IgE antibody to Trichophyton RAST units/ml

![Binding curve for IgE antibody to T. tonsurans. Serial twofold dilutions of the control serum were assayed in duplicate, mean (±), and range of duplicate values are indicated. Also presented is the mean ± SD for duplicate assays on sera from six nonallergic control subjects (o).]

For assays, 50 μl of Trichophyton Sepharose suspension was incubated with 100 μl of serum or serum diluted with 50% horse serum, and rotated overnight at room temperature in 12 mm by 75 mm plastic tubes. After washing three times with BBS, 125I-labeled goat anti-IgE (~100,000 cpm) was added in 200 μl of 50% horse serum and again rotated overnight at room temperature. Finally, the Sepharose was washed three times and counted in a Micromedic (Micromedic Systems, Horsham, Pa.) 4-channel automatic gamma counter. Six sera from nonallergic control subjects were assayed in duplicate in parallel with each assay, and values >2 SD greater than the mean for the control sera were regarded as positive. However, in assaying sera from allergic patients, it became evident that nonspecific binding was higher with sera from patients with high total IgE. Background binding was assessed in a series of skin test negative patients; background was 0.48 ± 0.07% for 28 sera with <100 IU of IgE per milliliter; 0.66 ± 0.18% for 30 sera with 100 to 1000 IU of IgE per milliliter, and 1.58 ± 0.95% for nine sera with >1000 IU of IgE per milliliter. All sera were assayed for total IgE, and relevant background was subtracted before calculating results.

**Paper disc RAST**

Cellulose discs were made from Whatman No. 1 filter paper and activated with CNBr with standard techniques. Discs were coated individually with Trichophyton extract diluted 1:20 in 0.1 mol/L NaHCO₃, 100 μl per disc, overnight. Dermatophagoides pteronyssinus RAST was carried out as described previously.

**RESULTS**

**RAST for IgE antibody to Trichophyton**

Initial experiments with Trichophyton extract linked to cellulose discs produced very poor binding, maximum binding 4% of added radioactivity. These results were improved either by extensive dialysis of extracts or by taking a 60% to 90% SAS precipitate (Table I). However, with dialyzed extract linked to Sepharose 4B, we obtained better binding (up to 30% of added anti-IgE bound) (Table I). With up to 3% bovine serum albumin in the RAST diluent, the nonspecific background was very high (up to 2.5%). Marked reduction in background was obtained by use of 50% horse serum as diluent in all phases of the assay. Nonetheless, background was higher for sera with high total IgE, and relevant background values were subtracted (see Methods). Both binding phases of the assay were found to require rotation of the tubes, and the assay elicited most consistent results with prolonged incubation periods, i.e., greater than or equal to 12 hours. In preliminary experiments with both cellulose discs and Sepharose, we found better binding to T. tonsurans and to Trichophyton mixture than to T. rubrum (these results are presented in part in Table I). One hundred sera were assayed both against T. tonsurans and Trichophyton mixture on Sepharose. The results were closely comparable with an overall correlation coefficient of r = 0.96 (p < 0.001) (data not presented). For the routine assay, T. tonsurans on Sepharose was used, and quantitation was obtained by assaying serial twofold dilutions of a strong positive serum that was arbitrarily allotted 1000 RAST units of IgE antibody to T. tonsurans (Fig. 1). Each test serum was assayed at two dilutions. Results were interpolated from the control curve, and mean intra-assay variation for the two dilutions was ± 16% of the mean. Repeated assays on 30 sera demonstrated a coefficient of variation of 27%.

Several approaches were used to assess the specificity of the assay. The results with different species of Trichophyton suggest that there is extensive cross-reactivity among them. All the sera were assayed for IgE antibody to D. pteronyssinus, and these results demonstrated no correlation with Trichophyton results (Fig. 2). We also examined skin test records on 45 of the patients. Although many of the patients had other positive skin tests, there was no consistent pattern. In particular 16/24 of the patients with strong positive skin tests to Trichophyton had no other com-
IgE antibodies to *Trichophyton*

**FIG. 2.** IgE antibody to *Trichophyton* and IgE antibody to *D. farinae* assayed in 106 sera. Fifty-five sera that demonstrated no IgE antibody to either allergen are indicated (0 x 55), below the lower limit for each assay. There was no correlation between the results for the two assays (r = -0.16) (p > 0.2).

parable (i.e., 2+) skin test to the other fungi tested. These results suggest that positive skin tests to other fungi are likely to represent concordant sensitivity rather than cross-reactivity. Finally, we assayed 61 sera for IgE antibody to *Alternaria*. Five sera in that group were positive for *Alternaria* alone, three sera to *Trichophyton* alone, and four sera were positive to both. Again the results suggest that the examples of dual sensitivity are likely to represent concordant sensitivity rather than cross-reactivity.

**IgE antibodies to *Trichophyton* in sera from patients with angioedema, urticaria, asthma, or rhinitis**

Among 27 patients with chronic urticaria and/or recurrent angioedema, we found eight patients with strong positive skin tests to *Trichophyton* and six additional patients with weaker skin reactions. Nine of these patients had serum IgE antibody to *Trichophyton*. Eight had strong positive skin tests plus one who demonstrated a 6 by 6 mm wheal (Fig. 3). Four of these patients had no other strong positive skin tests. The others demonstrated various positive skin tests including mite (two), ragweed (one), wheat (one), and Cladosporium (Hormodendrum) (one).

Strong positive skin tests to *Trichophyton* were not very common among patients with asthma. However, we have identified 15 patients with 2+ skin tests of approximately 120 who were skin tested. In sera from 12 of those 15 patients, we found detectable serum IgE antibody to *Trichophyton* with values ranging from 10 to 500 RAST units per milliliter (Fig. 3). Of the 12 patients with asthma who had serum IgE antibody to *Trichophyton*, four also had IgE antibody to *D. pteronyssinus*, whereas eight patients had none. In addition to clinic patients, we assayed sera from 61 patients with asthma observed in the emergency room and elicited positive results from seven sera (range 2.3 to 35 *Trichophyton* RAST units per milliliter). However, of these seven sera, six had multiple IgE antibodies to other allergens as assessed by RAST (data not presented). Indeed, the one patient whose serum contained IgE antibody to *Trichophyton* (35 U/µl) but was otherwise negative had already been identified as *Trichophyton* sensitive during a clinic visit and was one of the patients included in Fig. 3.

A strong positive (2+) immediate skin test was obtained in five of thirty-four individuals who worked in the hospital and were skin tested with *Trichophyton*. Of these individuals, four had detectable serum IgE antibody to *Trichophyton* (Fig. 3). All four of the control subjects who had IgE antibody to *Trichophyton* had had chronic fungal infection of the feet requiring local antifungal treatment, but none of them had had urticaria or asthma.
Traditionally, it was recognized that the different species of *Trichophyton* cross-reacted extensively but that they did not cross-react with other fungi. We obtained similar in vitro results with either *T. tonsurans*, *T. rubrum*, or mixed *Trichophyton* extract. Furthermore, our skin test results revealed no consistent pattern of sensitivity to other fungi in patients with positive skin tests to *Trichophyton* mixture. Although we have not carried out cross-absorption studies on sera from the minority of patients who were doubly sensitive, we think that these cases represent concordant sensitivity rather than cross-reactivity. In contrast, some patients who have strongly positive skin tests to other fungi demonstrate weak positivity to *Trichophyton*. In general, these weak reactions do not correlate with RAST results (Fig. 3). Although some of the patients who had positive RAST to *Trichophyton* were highly atopic, most were not. We have identified four patients with urticaria or angioedema and four with asthma who had no other positive skin tests. The mixed *Trichophyton* extract used here is the same extract that is widely used for testing for delayed hypersensitivity. Many of the patients and control subjects who did not demonstrate immediate skin tests developed delayed responses at 24 and 48 hours.

The finding of immediate hypersensitivity to a dermatophyte may simply reflect the widespread presence of fungal infections and their chronicity. *T. rubrum* and *T. mentagrophytes* are commonly implicated in foot, groin, and nail infections in our area, whereas *T. tonsurans* is more commonly implicated in head infections of children. The presence of IgE antibody may influence both the growth of these fungi and the local symptoms, e.g., itching of the feet or groin. In contrast, in patients who present with other diseases, the question arises whether the combination of *Trichophyton* allergy and absorption of fungal antigens from the skin could play a role in the other disease. This view was strongly encouraged by Sulzberger and Wise in 1930 but has been largely ignored in textbooks recently. However, there are two more recent articles that report urticaria may be related to *Trichophyton* infection, and in at least one case...
urticaria responded to antifungal treatment. We have recently reported that four of the present patients who had recurrent attacks of angioedema and/or anaphylaxis improved with antifungal treatment. We are not aware of any recent discussion of the role of *Trichophyton* allergy in asthma. It appeared possible in six of the present cases that *Trichophyton* could have a causative role. Improvement of asthma in two cases has coincided with treatment of the fungal infection. Since immediate hypersensitivity to *Trichophyton* is not restricted to patients with any one disease and occurs not infrequently in normal individuals, the association itself does not provide evidence for a causal relationship. Probably the only convincing evidence would have to come by demonstrating that antifungal treatment caused a cure or marked improvement of the associated condition. We have only occasional cases at present that have been treated in an uncontrolled fashion. We believe that our results establish the usefulness of RAST assay for confirming IgE-mediated sensitivity to this antigen and should serve to stimulate renewed interest in this area.

We thank the staff of the emergency room for help with collecting sera from patients with asthma, Gail Rose for excellent technical help. Nancy Malone for preparing the manuscript, and Hollister-Stier, Ltd. for generous supplies of *Trichophyton* extracts and culture.

REFERENCES


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